Carcinomatous meningitis: antibody-guided therapy with I-131 HMFG1

R P Moseley, J C Benjamin, R D Ashpole, N M Sullivan, J A Bullimore, H B Coakham, J T Kemshhead

Abstract

Seven patients with carcinomatous meningitis were administered intrathecal I-131 labelled monoclonal antibody HMFG1. Clinical responses were seen in two patients, with a long term survivor at 32 months. Aseptic meningitis occurred in 4/7 patients, but more serious toxicity was observed in the form of seizures (2/7 patients) and myelosuppression (3/7 patients). Partial obliteration of the subarachnoid space was identified as a potential problem in patients with advanced disease.

The development of monoclonal antibodies defining tumour associated antigens has significantly enhanced the field of diagnostic histopathology. As a natural extension of their success as immunocytochemical reagents, extensive investigation has explored their potential as vectors for targeting radioisotopes, toxins or drugs to tumours. In animal xenograft models, specific targeting of radioimmunoconjugates to tumour sites has been demonstrated. Therapeutic effect has also been observed in these models as evidenced by shrinkage and even cure of tumours.

In contrast to animal studies, the intravenous administration of monoclonal antibody immunoconjugates into patients with tumours has been consistently disappointing. In most studies, the percentage of the total injected dose of immunoconjugate localising within tumour tissue has been extremely low (approximately 10-35% of the injected dose per gram of tumour) and this has been attributed to several factors. Firstly, the volume of distribution of these radiopharmaceuticals in humans is several orders of magnitude greater than in laboratory animals and this dilutional effect clearly reduces the concentration of immunoconjugate at tumour sites.

Secondly, the penetration of immunoconjugate into tumours may be restricted by limited permeability of these macromolecules across lipid membrane blood/tumour barriers. Other contributing factors cited have included the presence of tumour-shed antigens, and the sequestration of blood pool immunoconjugate within abdominal viscera. Many of these problems are circumvented by the direct instillation of these reagents into body cavities where tumour cells are directly exposed to high concentration of immunoconjugate. These techniques were pioneered in the coelomic cavities and subsequently evaluated in the subarachnoid space of patients with neoplastic meningitis. Experience to date with intracavitary monoclonal antibody radioimmunotherapy has largely been with iodine-131 and yttrium-90. These predominantly beta-emitting radioisotopes deliver short range energy to micrometastases, whilst sparing normal body tissues. Neoplastic meningitis is characterised by the existence of thin sheets of leptomeningeal tumour and free floating malignant cell clusters within the subarachnoid space. It is therefore theoretically amenable to specifically targeted short range radiation delivered via the CSF pathways. In preliminary studies of antibody guided radiation in neoplastic meningitis, demonstrable therapeutic responses were seen in patients with relatively radio-sensitive neuro-ectodermal and lymphoid tumours. The efficacy of this method for the treatment of neoplastic meningitis secondary to carcinoma (carcinomatous meningitis) is also currently being evaluated and we report here the results obtained with seven patients treated by intrathecal administration of I-131 labelled monoclonal antibody HMFG1.

Methods

Antibody

The antibody HMFG1 was developed against the human milk fat globule and binds to an epitope on the high molecular weight glycoprotein (trypomorphic epithelial mucin, PEM). Tissue distribution of this antigen is limited to a variety of normal and neoplastic derivatives of glandular epithelium. Purified HMFG1 antibody (Unipath) in phosphate buffered saline was radiolabelled with the stationary phase chloramide iodogen to a specific activity of 5–10 uCi/ug of protein. Structural integrity of radioimmunoconjugates was assessed by trichloroacetic acid (TCA) precipitation and gel filtration chromatography with Sephacryl S300. Biological function of immunoconjugates was estimated by determination of the immunoreactive fraction by radiobinding assay at antigen excess. All radiopharmaceuticals were sterilised by millipore filtration before administration to patients.

Patient selection and management

Seven patients with carcinomatous meningitis were studied. Patients included in this study had either failed an adequate trial of conventional therapy or had been referred directly
following diagnosis. Patients and their relatives were fully informed of the experimental nature of this work and provided their written consent. The study was also approved by the Frenchay Hospital Ethical Committee. In all patients, a full clinical evaluation of disease was performed, supported by haematological and biochemical assessment. Cranial CT scanning was performed to exclude intracranial mass lesions. Pan-myelography and CT cisternography were subsequently employed to ascertain the patency of the CSF pathways. Immunocytochemistry of CSF was performed firstly to confirm the diagnosis of carcinomatous meningitis and secondly to demonstrate the presence of HMFG1 defined antigen (PEM) on malignant cells. In all patients, the diagnosis of carcinomatous meningitis was also confirmed by the demonstration of soluble PEM antigen in CSF.\textsuperscript{25}

Thyroid blockade was immediately started in all patients and continued for approximately one month following administration of immunon conjugate. Glucocorticoids (dexamethasone 1 mg three times daily) were also administered in an attempt to suppress aseptic meningitis which was seen in earlier patients following intrathecal administration of immunon conjugate.\textsuperscript{18} A single intrathecal administration of immunon conjugate was performed via an Ommaya reservoir ventriculoscopy following an isovolumetric withdrawal of CSF. Administered radioactivity dosage in patients 1–7 was 55, 55, 58, 56, 30, 60, and 54 mCi respectively. Patients were nursed in radioprotective facilities until levels of radioactivity had fallen sufficiently to allow conventional nursing care.

\textbf{Imaging}

Immunoscintigraphy of the neuraxis was performed with a General Electric MaxiCamera 400T equipped with a high energy parallel-hole collimator and linked to a DEC PDP 11/34 computer with gamma 11 software. Patients were scanned as soon and as often as their clinical condition permitted. In cases 2 and 3, clinical circumstances prevented acquisition of scanning biodistribution data.

\textbf{Evaluation of clinical response}

Patients were evaluated for response if they had received neither chemotherapy nor radiotherapy to all evaluable sites within the preceding six weeks. These conditions were waived if the patients had clear evidence of disease progression in the intervening period.

All patients in this study were initially considered clinically evaluable by these criteria. Response was assessed at three-monthly intervals by clinical examination, cranial CT scanning and cytological examination of CSF. In addition, estimation of PEM in serum and CSF at three months following therapy was performed in cases 1, 3, 4, 5, and 7.

\textbf{Results}

Seven patients with carcinomatous meningitis were studied, with a range of primary tumour sites (table 1). The age range was 36–60 years and there were three males and four females. The immunoreactivity of administered conjugates exceeded 50% in all patients.

\textbf{Immunoscintigraphy}

I-131 activity was observed within the cerebral ventricles and craniospinal subarachnoid space in all patients who had gamma scintigraphy. However, neuraxial distribution of radionuclide was always irregular and incomplete. Paucity of radionuclide was commonly observed over the cerebral hemisphere convexities or within the basal subarachnoid cisterns (fig 1). Sites of increased neuraxial isotope uptake were frequently demonstrated at clinically suspected and/or radiologically demonstrated sites of disease (tables 2 and 3).

\textbf{Clinical response to therapy}

Pre-treatment status and clinical responses in all seven patients are shown in tables 4 and 5. Clinical responses were assessed in terms of neurological response, CSF response and survival. Neurological response was categorised as complete (CR) if there was total resolution of symptoms and signs. Partial response

\begin{table*}
\centering
\begin{tabular}{|c|c|c|c|}
\hline
\textbf{Patient} & \textbf{Age in years} & \textbf{Sex} & \textbf{Primary tumour} & \textbf{Sites of clinically apparent extraneuraxial disease} \\
\hline
1 & 48 & F & Ovarian carcinoma & — \\
2 & 60 & F & Bladder carcinoma & Liver \\
3 & 44 & F & Breast carcinoma & — \\
4 & 57 & M & Large cell lung carcinoma & Lung \\
5 & 41 & M & Lung adenocarcinoma & Lung \\
6 & 36 & F & Breast carcinoma & — \\
7 & 50 & M & Large cell lung carcinoma & Lung \\
\hline
\end{tabular}
\caption{Clinical parameters of patients with carcinomatous meningitis}
\end{table*}

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Cranial scintigram of case 5 at eight days revealing paucity of isotope activity over left cerebral hemisphere convexity. Necropsy studies revealed bilateral hemisphere disease involving the leptomeninges.}
\end{figure}
(PR) implied some improvement in neurological symptoms and signs. When neurological symptoms/signs remained stable or deteriorated, this was categorised as no response (NR). CSF response was assessed on the presence or absence of malignant cells seen on cytological examination. A complete response (CR) indicated complete eradication of malignant cells from two daily consecutive CSF preparations. A partial response (PR) indicated a sustained reduction of malignant cell count in two daily consecutive CSF preparations. When malignant cell counts remained unaltered or increased, it was considered that no response (NR) to therapy had been demonstrated. In cases 1, 3, 4, 5, and 7, an additional parameter of CSF response was obtained by estimation of HMFG1 (PEM) mucin in serum and CSF before and at three months following therapy.

Patient 2 died suddenly at 72 hours and was therefore clinically irrevocable for response. The remaining six patients were considered clinically evaluable for response. Neurological responses were seen in 2/6 patients (cases 1 and 3). CSF responses were seen in 1/6 patients (case 3). The latter patient remains in complete remission at 36 months, having received no additional treatment. In 4/5 patients, CSF levels of HMFG1 (PEM) mucin increased following therapy (cases 1, 4, 5, and 7), indicating progressive disease within the central nervous system (fig 2). In case 3, a positive diagnostic assay result became negative after therapy. In this same group of patients with the exception of case 3, serum HMFG1 (PEM) mucin increased following therapy, indicating progressive systemic disease (fig 3). Serum levels of HMFG1 (PEM) mucin in case 3 remained approximately constant, indicating stable systemic disease.

Case report (patient 3) A 40 year old woman had simple mastectomy for breast carcinoma. Four years later, a cervical lymph node metastasis was excised with local irradiation. She was started on tamoxifen and had radiation oophorectomy. Several months later she developed ataxia, diplopia and facial weakness. Cranial CT was normal, but HMFG1-PEM and malignant cells were demonstrated in the CSF. Following intravenous chemotherapy and 3000 cGy of whole brain irradiation, she deteriorated with the development of a complete third nerve palsy, nystagmus, masseter wasting, facial numbness, deafness, bulbar paresis, saddle numbness and loss of deep tendon jerks. Instillation of intraventricular I-131 immunon conjugate was initially uneventful, but 10 days later she developed status epilepticus requiring ventilation. Following the control of seizures she was discharged home. At three months following discharge her neurological status remained unchanged, but lumbar CSF was free of malignant cells and HMFG1-PEM. She subsequently gained weight and improved neurologically. At 18 months following discharge, she had recovered facial sensation and normal gait. Deep tendon jerks had reappeared and saddle numbness had resolved. She remained with a partial third nerve palsy and deafness but was able to continue normal activities independently. At 36 months she remains in remission with absence of malignant cells and HMFG1-PEM in CSF.

Toxicity The commonest form of toxicity experienced was an aseptic meningitis manifesting as headaches, nausea, vomiting, nuchal rigidity and pyrexia. This was observed in cases 1, 4, 5, and 6, and in all instances resolved spontaneously over approximately 48 hours. The fatality of case 2 falls into the category of an acute toxic complication of therapy which occurred at approximately 72 hours after instillation of immunon conjugate. Necropsy studies revealed no obvious cause of her sudden death, but there was circumstantial evidence to suggest that this may have been due to unreported epileptic seizures. Epilepsy was part of her presenting clinical syndrome and 12 hours before her death she had been observed in a transient semi-comatose state lasting approximately 20 minutes. New onset epilepsy was subsequently seen in case 3 at 10 days following administration of immunon conjugate. Three patients developed bone marrow suppression (cases 1, 3, and 6). In cases 1 and 3, recovery of pancytopenia was spontaneous. The nadir in white cell count occurred at week 5 in patient 1 (WCC: 3.1 x 10^9/lit) and week 4 in patient 3 (WCC: 2.1 x 10^9/lit). A commensurate fall in platelet count was noted, reaching 86 x 10^9/lit and 76 x 10^9/lit respectively. In both patients a return to normal blood parameters was observed by week 9. In patient 6, the development of a severe pancytopenia at four weeks (WCC: 8.9 x 10^9/lit; platelets 6 x 10^10/lit) was accompanied by a gram negative septicemia requiring parenteral antibiotics.

### Table 2 Intrathecal therapy: diagnostic radiology

<table>
<thead>
<tr>
<th>Patient</th>
<th>CT/myelography</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Excess CSF over cerebral hemisphere and nodular leptomeningeal tumour involving cauda equina</td>
</tr>
<tr>
<td>2</td>
<td>Hydrocephalus</td>
</tr>
<tr>
<td>3</td>
<td>Normal</td>
</tr>
<tr>
<td>4</td>
<td>Hydrocephalus, nodular leptomeningeal tumour involving cauda equina</td>
</tr>
<tr>
<td>5</td>
<td>Multiple low density areas seen over cerebral hemisphere and nodular leptomeningeal tumour involving cauda equina</td>
</tr>
<tr>
<td>6</td>
<td>Hydrocephalus, diffuse contrast enhancement over left cerebral hemisphere convexity</td>
</tr>
<tr>
<td>7</td>
<td>Nodular leptomeningeal tumour involving cauda equina</td>
</tr>
</tbody>
</table>

### Table 3 Intrathecal therapy: immunoscintigraphy

<table>
<thead>
<tr>
<th>Patient</th>
<th>Extent of neovascular distribution</th>
<th>Sites of increased neovascular distribution</th>
<th>Sites of extraneovascular distribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Incomplete</td>
<td>Cranial vertex</td>
<td>Thyroid</td>
</tr>
<tr>
<td>2</td>
<td></td>
<td>Lumbar/sacral spine</td>
<td>Abdominal Foci</td>
</tr>
<tr>
<td>3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Incomplete</td>
<td>Lumbar/sacral spine</td>
<td>Spleen</td>
</tr>
<tr>
<td>5</td>
<td>Incomplete</td>
<td>Left hemisphere convexity</td>
<td>Spleen</td>
</tr>
<tr>
<td>6</td>
<td>Incomplete</td>
<td>Thoracic spine</td>
<td>Diffuse thoracic</td>
</tr>
<tr>
<td>7</td>
<td>Incomplete</td>
<td>Lumbar/sacral spine</td>
<td>Diffuse thoracic</td>
</tr>
</tbody>
</table>

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and transfusion of blood products. It was noteworthy that a peripheral blood count before immunoconjugate administration revealed thrombocytopenia (platelets 86 x 10^9/lit) which may have reflected tumour infiltration of bone marrow. She died at six weeks following therapy and no necropsy was performed.

**Discussion**

Carcinomatous meningitis may complicate any form of systemic carcinoma, but the most commonly reported primary tumour sites include breast, lung, and gastrointestinal tract. The disorder is generally associated with a poor prognosis and response to therapy. Median survival with untreated disease is often reported as four to six weeks, although this may reflect delayed clinical diagnosis. The conventional approach to the management of this condition involves intrathecal chemotherapy combined with external beam irradiation to the site of major clinical involvement in the neuraxis. Reported clinical responses are variable, with median survivals ranging from two to 5-8 months. Survival and therapeutic response data appear consistently more favourable for breast carcinoma than lung carcinoma. Indeed, survivals of two to three years have been reported in occasional patients with neoplastic meningitis complicating breast carcinoma.

Although clinical responses have followed conventional therapeutic methods, the nonspecific nature of these modalities results in severe dose-limiting toxicity and consequent limitations in efficacy. As a possible solution to this therapeutic dilemma, several investigators have advocated the intrathecal instillation of colloidal radioactive gold or yttrium-DTPA. The short range beta-particle emissions from these radionuclides deposit their radiation energy within leptomeningeal tissues whilst sparing the underlying neural parenchyma. Although therapeutic success has been reported with this method, the occurrence of cauda equina neurotoxicity has limited enthusiasm for its use. It is hoped that linkage of short range emitting radionuclides to monoclonal antibodies may facilitate greater specificity of radiation delivery, with consequent reduction in neurotoxicity.

By our defined criteria, clinical responses were seen in 2/6 evaluable patients (cases 1 and 3). In case 3, a complete CSF response was observed as evidenced by the elimination of malignant cells and soluble HMFG1 (PEM) mucin. In cases 1, 4, 5, and 7, CSF HMFG1 (PEM) mucin levels increased over a three month period following therapy and malignant cells persisted. In the same patients, serum levels of HMFG1 (PEM) mucin also increased, indicating progressive systemic disease. It is conceivable that, in these patients, systemic tumour may have continued to seed the leptomeninges following therapy. In carcinoamatous meningitis, stable systemic disease may therefore be a prerequisite for the eradication of leptomeningeal tumour with a single instillation of immunoconjugate. Clinically demonstrated extraneuraxial disease may in itself be stable. In these circumstances intrathecal radioimmunotherapy may provide useful palliation of severely debilitating symptoms and sequential serum estimation of PEM may provide evidence to support the stability of systemic disease. Primary neuroectodermal

### Table 4 Intrathecal therapy: pretreatment status

<table>
<thead>
<tr>
<th>Patient</th>
<th>Prior therapy (CNS)</th>
<th>Disease status (CNS)</th>
<th>Clinical status</th>
<th>CSF cytology</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CS radiotherapy, IT chemotherapy</td>
<td>Progressive disease on treatment</td>
<td>Diplopia, paraparesis, urinary incontinence</td>
<td>+</td>
</tr>
<tr>
<td>2</td>
<td>VP shunt</td>
<td>New disease</td>
<td>Dementia, epilepsy, limb spasticity</td>
<td>+</td>
</tr>
<tr>
<td>3</td>
<td>Craniotomy</td>
<td>New disease</td>
<td>Confusion, multiple cranial nerve palsies</td>
<td>+</td>
</tr>
<tr>
<td>4</td>
<td>IT chemotherapy</td>
<td>Progressive disease on treatment</td>
<td>Paraparesis</td>
<td>+</td>
</tr>
<tr>
<td>5</td>
<td>VP shunt</td>
<td>New disease</td>
<td>Headache, backache, meningism</td>
<td>+</td>
</tr>
<tr>
<td>6</td>
<td>IT chemotherapy</td>
<td>Progressive disease on treatment</td>
<td>Nausea, vomiting, headache, meningism</td>
<td>+</td>
</tr>
<tr>
<td>7</td>
<td>Craniotomy</td>
<td>Progressive disease on treatment</td>
<td>Deafness, paraparesis, facial weakness and numbness</td>
<td>+</td>
</tr>
</tbody>
</table>

C/S, craniospinal; IT, intrathecal; VP, ventriculo-peritoneal.

### Table 5 Intrathecal therapy: clinical responses (November 1990)

<table>
<thead>
<tr>
<th>Patient</th>
<th>Neurological response</th>
<th>CSF response</th>
<th>Survival (months)</th>
<th>Current disease status (CNS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>PR</td>
<td>NR</td>
<td>7*</td>
<td>--</td>
</tr>
<tr>
<td>2</td>
<td>Toxic</td>
<td>Death</td>
<td>D</td>
<td>--</td>
</tr>
<tr>
<td>3</td>
<td>PR</td>
<td>CR</td>
<td>36</td>
<td>Remission</td>
</tr>
<tr>
<td>4</td>
<td>NR</td>
<td>NR</td>
<td>4*</td>
<td>--</td>
</tr>
<tr>
<td>5</td>
<td>NR</td>
<td>NR</td>
<td>3*</td>
<td>--</td>
</tr>
<tr>
<td>6</td>
<td>NR</td>
<td>NR</td>
<td>1.5*</td>
<td>--</td>
</tr>
<tr>
<td>7</td>
<td>NR</td>
<td>NR</td>
<td>5*</td>
<td>--</td>
</tr>
</tbody>
</table>

CR, complete response; PR, partial response; NR, no response. *Patient dead. D, patient died at 72 hours following injection.

![Figure 2](http://jnnp.bmj.com/) Estimation of CSF HMFG1 (PEM) mucin before and at three months following therapy.
Figure 3 Estimation of serum HMFG1 (PEM) mucin before and at three months following therapy.

Meningeal neoplasia may offer a more favourable pathology than that of carcinomatous meningitis as the former is generally more radiosensitive and disease is confined within the central nervous system and therefore theoretically accessible to intrathecal administered immunoconjugate.

In this study, acute toxicity in the form of an aseptic meningitis occurred in over 50% of patients. Similar toxicity has been reported following intrathecal administration of methotrexate, 46 cytosine arabinoside 46 and radioiodinated human serum albumin (RISA). 46 The reported incidence of aseptic meningitis following intrathecal RISA has been highly variable and this complication has been attributed to contaminating endotoxins. 47 Aseptic meningitis in our patients was unrelated to administered radioactivity dose and was observed intermittently. It is likely that the presence of contaminating pyrogens accounts for this complication, although radio-pharmaceuticals were not tested for their presence. Epilepsy occurred in two patients. We have considered this a toxic response to therapy, but of course seizures occur as a natural event in the disorder. 48 Delayed central nervous system toxicity has not been observed clinically in long term survivors, but a necropsy study in case 2 revealed extensive periventricular and brainstem sub-pial white matter oedema and astrocytic reaction. 48 Bone marrow suppression was seen in three patients, and appears to be the dose-limiting toxicity of this form of therapy.

Before dismissing these clinical results as unimpressive, one should consider the devastating nature of this disease. Many of these patients had advanced disease, unresponsive to previous therapy. The important observation here is that therapeutic responses have been demonstrated. As with other forms of oncolytic therapy, clinical results may be more favourable with minimal disease. As leptomeningeal disease progresses, partial obliteration of the subarachnoid space may prevent complete distribution of immunoconjugate throughout the CSF pathways and this clearly reduces the opportunity for therapeutic success. 49 Emphasis should therefore be placed on the diagnosis of minimal disease and on establishment of complete patency of the craniospinal subarachnoid space before therapy. The addition of Indium-111-DTPA ventriculography may complement CT myelography in this assessment, 46 and areas of reduced radionuclide distribution may be externally irradiated before intrathecal radioimmunotherapy.

In conclusion, we suggest that intrathecal instillation of monoclonal antibody radioimmunoconjugates may produce a worthwhile clinical response in carcinomatous meningitis, although this may be less marked than that observed in more radiosensitive lymphoid and primary neuroectodermal tumours. Modifications in the methodology may further enhance the therapeutic efficacy of the technique. Firstly, monoclonal antibodies defining alternative epithelial specific antigens might be employed with advantage. A theoretical concern with monoclonal antibody HMFG1 relates to the glycoprotein antigen (PEM) it defines. Polymorphic epithelial mucin (PEM) is liberated from the cell surface into CSF and this may reduce tumour cell targeting of immunoconjugates. 49 Further variations may include multiple doses, alternative radionuclides or the use of antibody cocktail mixtures to account for heterogeneity of tumour cell antigen expression. 90 We believe that intrathecal radioimmunotherapy should be pursued as an alternative method of palliative treatment for carcinomatous meningitis, as presently existing methods of treatment are clinically inadequate. The value of this technique will become more apparent when a larger experience has been accumulated.

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References

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